IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Appl. No. : 10/571.508 Confirmation No.: 7832

Applicant (s) : Liping Zhang et al.

Filed : March 10, 2006

TC/A.U. : 1625

Examiner : Bernard I Dentz

Title : A PROCESS FOR THE PRODUCTION OF ALKYLENE OXIDE

USING A GAS-PHASE PROMOTER SYSTEM

Docket No. : 63261A Customer No. : 00109

Declaration Under 37 C.F. R. § 1.132

The undersigned, Dr. Liping Zhang, citizen of the United States of America and residing in the State of Texas, United States of America declare and say that:

- I am a 1982 graduate of Tsing-hua University, Beijing, China with a Bachelor of Science degree in Chemical Engineering, a 1985 graduate of Tsing-hua University, Beijing, China with a Masters of Science degree in Chemical Engineering and a 1996 graduate of The Ohio State University, Columbus, Ohio, USA with a Doctor of Philosophy Degree in Chemical Engineering.
- 2. From April 1985 until April 1991, I was employed as an R&D Engineer at the Research Institute of Petroleum Processing, SINOPEC Co. in China. From September 1987 until December 1988 and then from May 1991 until September 1992, I was a Visiting Research Associate at The Ohio State University. From September 1992 through November 1996, I was a Graduate Research Associate with The Ohio State University.
- 3. In December 1996, I joined Union Carbide Corporation ("UCC") in South Charleston, West Virginia as a Senior Engineer. I held this position until March of 2000. At that time, I became a Project Scientist with UCC, and held this position until April, 2001, when UCC was acquired by The Dow Chemical Company ("TDCC"). After UCC became a wholly owned subsidiary of TDCC, I was named a Research Specialist. In March 2003, I was promoted to Senior Research Specialist; this is my role today. I

worked at the South Charleston site until 2008 when I relocated to TDCC's Freeport, Texas, location.

- 4. In my roles at UCC and TDCC, my responsibilities included leading the design, construction, and commissioning of the Ethylene Oxide ("EO") R&D ROTOBERTY® laboratory, the supervision of operation of ROTOBERTYs, autoclaves, pilot plant, and microreactor systems, conducting studies of catalyst formulation, catalyst preparation, catalyst characterization, and kinetics to improve the performance and fundamental understanding of EO catalysts. I also developed methods and experiments for EO catalyst characterization and kinetic studies, provided technical support for EO plants, and developed gas phase promoter strategies for EO plants.
- I am currently an inventor on one US published application in the field of ethylene oxide production.
- 6. I have read the above referenced patent application, the Office Action dated February 13, 2009, and the prior art references, Thorsteinson et al., EP 425020 B1 ("Thorsteinson I"), Thorsteinson et al., EP 480537 ("Thorsteinson II"); and Hayden et al., US 5.387.751 cited by the Examiner.
- 7. Thorsteinson I does not disclose any catalysts which have produced 1.1 kilo metric tons of ethylene oxide per cubic meter of catalyst. In addition, I have determined the N*/Z* ratios for the relevant examples of Thorsteinson I, under the variety of conditions set forth in Thorsteinson I. The N*/Z* ratios for Thorsteinson I range from 0.18 to 2.99. Only three of 34 experiments use an N*/Z* ratio of between 0.4 to 1.0. Among these three experiments, i.e., Examples 28, 32, and 33, Example 33 had both N* (38) and Z* (80) values above the respective 20 and 40 values, as set forth in independent claim 1. Both Examples 28 and 33 are conducted under CO₂ concentrations of 3.0 mole %. There is no experiment in which all variables were constant except for N*/Z*. Thorsteinson I contains no teaching of the benefit of the claimed N*/Z* ratio for fresh catalysts, let alone for catalysts which have produced the claimed amount of ethylene oxide. See Table 1, attached to this declaration, for a list of the experimental conditions.

Examples 1-6 and 7-12

Examples 13-15

N*/Z*: 0.18

Examples 16-21

N*/Z* O₂ Condition: 0.26 N*/Z* Air Condition: 0.22

Examples 22-27

N*/Z* 0.26

Examples 28-31

N*/Z* Condition I: 0.93

N*/Z* Condition II: 0.27

N*/Z* Condition III: 2.99

N*/Z* Condition IV: 1.36

Examples 32-34

N*/Z* Air Condition: 0.51, where inlet C2H4% = 12 N*/Z* O₂ Condition I: 0.47, where N* =37.7 and Z* = 80.0 N*/Z* O₂ Condition II: 0.38

8. Similarly, Thorsteinson II does not disclose any catalysts which have produced 1.1 kilo metric tons of ethylene oxide per cubic meter of catalyst. In addition, I have determined the N*/Z* ratios for the relevant examples of Thorsteinson II. The N*/Z* ratios range from .20 to .48. Only 5 of 13 experiments in Thorsteinson II use an N*/Z* ratio between 0.4 and 1.0. For Catalyst Nos. 7, 8, and 9, the Z* value is 66.7, far above the value of 40 set forth in independent claim 1. There is no experiment in which all variables were constant except for N*/Z*. Thorsteinson II contains no teaching of the benefit of the claimed N*/Z* ratio for fresh catalysts, let alone for catalysts which have

produced the claimed amount of ethylene oxide. Instead, analysis of the data in Table 2 of Thorsteinson II shows that for younger catalysts, a lower N*/Z* ratio provides better performance. Compare Catalyst 4 at 20 days, having an N*/Z* ratio of 0.35, with an efficiency of 85.8%, % EO of 1.98, and temperature of 255C with Catalyst Nos. 1-3, which have an N*/Z* ratio of greater than 0.4. The increase in the N*/Z* ratio is coupled with a decrease in efficiency, or decrease in %EO as compared to Catalyst No. 4.

Catalyst No.	N*	Z*	N*/Z*
1	6.9	14.7	0.47
1	7.2	15.0	0.48
1	7.1	16.3	0.44
2	7.2	15.7	0.46
3	9.2	20.7	0.45
4	6.9	20.0	0.35
5	4.3	13.3	0.33
6	13.0	66.7	0.20
7	13.0	66.7	0.20
8	13.0	66.7	0.20
9	13.0	66.7	0.20
10, Condition I	4.3	16.7	0.26
10, Condition II	10.4	33.3	0.31

9. Similarly, Hayden does not disclose any catalysts which have produced 1.1 kilo metric tons of ethylene oxide per cubic meter of catalyst. In addition, I have determined the N*/Z* ratios for the relevant examples, under the varying conditions of Hayden. Where greater than zero, the N*/Z* ratios for Hayden range from 0.0006 to 0.099. Thus, none are within the claimed range. In Examples 13, 20, 29, 34 and 15, Z* is in excess of 66.7, far above the value of 40 set forth in independent claim 1. Hayden

contains no teaching of the benefit of the claimed N*/Z* ratio for fresh catalysts, let alone for catalysts which have produced the claimed amount of ethylene oxide. See Table 2, attached to this declaration, for the experimental conditions.

10. The following experiment demonstrates that a catalyst which has generated more than 1.1 kilo metric tons of ethylene oxide per m3 of catalyst benefits from operating within the claimed N*/Z* ratio.

Tubular fixed bed reactors contain a mixture of catalysts which have been used to produce ethylene oxide. The N*/Z* ratio was decreased from greater than 1 to 0.6. Catalyst efficiency (selectivity) improved by more than 1% without the loss of catalyst activity.

Properties of fresh catalysts in the reactor:

	Average	High	Low
Silver content (wt% by titration)	>30	>30	>30
Promoter	Potassium	Potassium	Potassium

Reactor inlet conditions:

Pressure (kPa, absolute)	2270 to 2310
Gas hourly space velocity (h-1)	6100 to 6900
Ethylene concentration (mol %)	21 to 27.5
Ethane concentration (mol %)	0 to 0.03
Oxygen concentration (mol %)	3 to 8.7
Carbon dioxide concentration (mol %)	0.2 to 0.5
Water concentration (mol %)	0.8 to 1.2
Ethylene dichloride (ppm)	0.3 to 1.1
Vinyl chloride (ppm)	0.5 to 2.0
Ethyl chloride (ppm)	0.0 to 0.03
Ammonia feed (ppm)	3.7 to 12.8
Total Nox (ppm) (estimated value based on	0.4 to 1.3
material balance)	
Balance gas: methane + nitrogen + argon	

The work rate (also referred to as the catalyst ethylene oxide loading) was between 150 to 270 (kgEO/h/m3 catalyst). The reactor coolant temperature ranged from 218 to 240°C.

The attached plots contain the "normalized" relative efficiency and "normalized" relative coolant temperature vs. cumulative EO loading. The normalized efficiency values were obtained by using the catalyst kinetic correlations to normalize the raw efficiency values to a reference work-rate of 238 kgEO/h/m3. The normalized efficiency values were then converted to the relative efficiency values by subtracting a constant reference efficiency number to convenient the comparison. The normalized coolant temperature values were obtained by using the catalyst kinetic correlations to normalize the raw coolant temperature values to a reference 238 kgEO/h/m3 work-rate and a reference 8.5% (mole) inlet oxygen concentration. The normalized coolant temperature values were then converted to the relative coolant temperature values by subtracting the same reference coolant temperature number to convenient the comparison.

Fig. 1 depicts N*/Z* on the left y-axis, N* or Z* on the right y-axis, and cumulative catalyst loading (kt EO/m3) along the x-axis.

Fig. 2 depicts N^*/Z^* on the left y-axis, relative efficiency (%) on the right y-axis and cumulative catalyst loading (kt EO/m3) along the x-axis.

Fig. 3 depicts Fig. 2 depicts N*/Z* on the left y-axis, normalized coolant temperature (C) on the right y-axis and cumulative catalyst loading (kt EO/m3) along the x-axis.

The figures demonstrate that operating the process at an N^*/Z^* ratio of between 0.4 to 1.0 for a catalyst which has produced in excess of 1.1 kilo metric tons of ethylene oxide per m3 of catalyst provides a selectivity increase but does not affect activity. Further, the figures demonstrate that the benefit of operating at the claimed N^*/Z^* ratio range is greater for the catalyst which has produced in excess of 1.1 kilo metric tons of ethylene oxide per m3 of catalyst than the benefit to a catalyst which has produced less than that volume of ethylene oxide.

Data between 0.42 to 0.51 ktEO/m3 cumulative catalyst loading were not plotted due to some uncertainty on measurements accuracy.

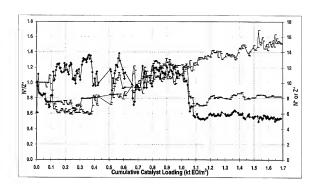


Fig. 2

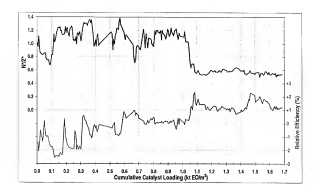
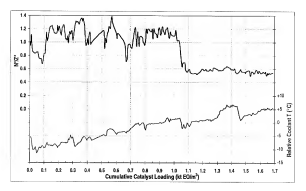


Fig. 3



11. I declare further that all statements made herein of my knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title

18 of the United States Code, and that willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Respectfully submitted,

Lining Zhang

Table 1

Examples 1-6 and 7-12																	
		۵		§	ż	걸	C2H4	C2H6	.2	.Z/.N	200	8	을 들	Day	-	GHSV	
		psig	кРа	wdd		mdd	%	%			%	%	%		ပ္	7	
Example 1	Condition I	275	1997	2	4.3	5	30.0	0.0	16.7	0.26	0	8.0	0.90	50	240	8000	
	Condition II	275	1997	55	13.0	20	30.0	0.3	33.3	0.39	3.0	8.0	0.17	20	270	8000	
Example 2	Condition I	275	1997	5	4.3	5	30.0	0.0	16.7	0.26	0	8.0	1.60	56	240	8000	
	Condition II	275	1997	15	13.0	50	30.0	6.0	33.3	0.39	3.0	8.0	0.16	56	270	8000	
Example 3	Condition I	275	1997	2	4.3	2	30.0	0.0	16.7	0.26	o	8.0	1.60	23	240	8000	
	Condition II	275	1997	15	13.0	50	30.0	6.0	33.3	0.39	3.0	8.0	0.40	23	270	8000	
Example 4	Condition I	275	1997	2	4.3	2	30.0	0.0	16.7	0.26	0	8.0	1.60	32	240	8000	
-	Condition II	275	1997	12	13.0	20	30.0	0.3	33.3	66.0	3.0	8.0	29.0	32	270	8000	
Example 5	Condition I	275	1997	2	4.3	2	30.0	0.0	16.7	0.26	0	8.0	1.50	48	240	8000	
	Condition II	275	1997	15	13.0	8	30.0	0.3	33.3	0.39	3.0	8.0	0.83	48	270	8000	
Example 6.	Condition 1	275	1997	2	4.3	2	30.0	0.0	16.7	0.26	0	8.0	1.60	40	240	8000	
-	Condition II	275	1997	15	13.0	50	30.0	0.3	33.3	0.39	3.0	8.0	0.80	40	270	8000	
Example 7	Condition I	275	1997	2	4.3	S.	30.0	0.0	16.7	0.26	0	8.0	1.69	2	240	8000	
-	Condition II	275	1997	15	13.0	50	30.0	0.3	33.3	0.39	3.0	8.0	1.71	6	270	8000	
Example 8	Condition 1	275	1997	2	4.3	2	30.0	0.0	16.7	0.26	0	8.0	1.82	2	240	8000	
	Condition II	275	1997	15	13.0	50	30.0	0.3	33.3	0.39	3.0	8.0	1.97	6	270	8000	
Example 9	Condition I	275	1997	5	4.3	2	30.0	0.0	16.7	0.26	0	8.0	1.85	-	240	8000	
	Condition II	275	1997	15	13.0	50	30.0	0.3	33.3	0.39	3.0	8.0	2.00	12	270	8000	
Example 10	Condition I	275	1997	2	4.3	വ	30.0	0.0	16.7	0.26	0	8.0	2.03		240	8000	
	Condition II	275	1997	15	13.0	50	30.0	0.3	33.3	0.39	3.0	8.0	2.07	4	270	8000	
Example 11	Condition I	275	1997	5	4.3	5	30.0	0.0	16.7	0.26	0	8.0	1.54	10	240	8000	
	Condition II	275	1997	15	13.0	20	30.0	0.3	33.3	0.39	3.0	8.0	1.61	13	270	8000	
Example 12	Condition I	275	1997	5	4.3	2	30.0	0.0	16.7	0.26	0	8.0	2.12	7	240	8000	
	Condition II	275	1997	15	13.0	20	30.0	0.3	33,3	0.39	3,0	8.0	1.96	6	270	8000	

	GHSV	ī	8000	8000	8000
	-	ပွ	236	236	236
	Day				
EO	ont	%	1.30	1.45	1.10
	8	%	8.0	8.0	8.0
	200	%	3.0	3.0	3.0
	,Z,		0.18	0.18	0.18
	ž		33.3	33.3	33.3
	C2H6	%	0.0	0.0	0.0
	C2H4	80	30.0	30.0	30.0
	딥	mdd	10	10	9
	ż		6.1	6.1	6.1
	ş	mdd		2	7
		кРа	1997	1997	1997
	۵	bsid	275	275	275
			Example 13	Example 14	Example 15

Examples 16-21

GHSV

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> 8 % 8 % 0.0 4.0 4.0 4.0 4.0

.Z/.N

Ň C2H6 88 0.3 0.3 0.3

C2H4 30.0 8.0 8.0

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> ķРа 1997

psig 275 275 275 275 275

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4.3 4.3 6.9 6.9

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8000 8000 0008

1.10

0.8 0.0 0.0 6.0

0.26 0.22 0.22 0.22

16.7 31.6 31.6 31.6 31.6

0.45 1.90 0.40 0.30 06.0

O₂ Condition O₂ Condition Example 16 Example 18 Example 17 Example 19

Air Condition Air Condition Air Condition Air Condition

1997 1997 1997

Example 20 Example 21

Examples 22-27

Example 22
Example 23
Example 25
Example 24
Example 26
Example 26
Example 26

	_				_		_	
2	SHSV	Æ	8000	8000	8000	8000	8000	8000
ŀ		ပ္	223	218	220	220	220	220
)							
요 .	out	%	1.04	1.51	1.30	1.50	1.12	1.95
8	SS	%	8	8	8	8	8	80
č	SSS	%	0	0	0	3	3	9
į	N./Z.		0.26	0.26	0.26	0.26	0.26	0.26
i	.7		16.7	16.7	16.7	16.7	16.7	16.7
	CZHP	%	0.0	0.0	0.0	0.0	0.0	0.0
	SZH4	%	30.0	30.0	30.0	30.0	30.0	30.0
-	2	mdd	5	5	5	2	5	5
	ż		4.3	4.3	4.3	4.3	4.3	4.3
	2	шdd	2	5	2	2	5	5
		kPa	1997	1997	1997	1997	1997	1997
	a.	psig	275	275	275	275	275	275

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Example 28
Example 29
Example 30
Example 31

	_	၁့	250	260	243	258
	击	%	88	88	87	98
ЕО	ont	%	1.00	0.35	2.22	2.30
	05	%	7.5	8.0	8.5	8.5
	202	%	3.0	1.0	3.0	5.0
	N*/Z*		0.93	0.27	2.99	1.36
	,Z		15.0	52.3	12.0	29.2
	C2H6	%	0.3	0.2	0.2	0.5
	C2H4	%	20.0	24.0	30.0	28.0
	ECL	ррт	7.5	23.0	6.0	14.0
	ż		13.9	14.1	35.8	39.7
	80	mdd	16	20	38	25
		kPa	1997	1618	2170	1756
	Д	psig	275	220	300	240

GHSV h-1 6200 4700 3600 3300 GHSV 5300 3600 6600

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82.0 89.0

8 8 8 8 3.0 % .Z.N 0.47 20.8 80.0 15.0 Ň

0.2

25 75 30.0 PB 45 8 10.6 37.7 5.7 ž

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Examples 32-34

Exampl Exampl Exampl

		psig	кРа	mdd
ple 32	Air Condition	220	1618	15
ple 33	O2 Condition	300	2170	4
ple 34	O2 Condition	300	2170	9

Table 2

Example No.	P (psia)	C2H4 (%)	C2H6%	VCL ppm	NO ppm	*N	*Z	*Z/*N
2	240	30	0	10	3	2.16	33.3	90.0
4	20	30	0	10	5	0.30	33.3	600.0
6	240	30	0	10	2	1.44	33.3	0.043
13	19.7	30	0	20	10	0.59	2.99	600.0

0.043	0.058	0.001	0.001	
33.3	18.7	66.7	2.99	
1.4	1.08	90:0	90'0	
2	1.5	1	1	
01	17	20	20	
0	9:0	0	0	
30	31	30	30	
240	240	19.7	19.7	
25	26	29	34	

0.008

2.99

0.53

12

20

0

30

14.7

20

19.7	30	0	20	-	90.0	2.99	0.001

*Z/*N 0.043 0.033

ž2

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NO ppm

VCL ppm

C2H6%

C2H4 (%)

P (psia)

Example Nos. 10

and 11

33.3 21.7 23.3 25.0 26.7 30.0 33.3 23.3 30.0 33.3 33.3

1.44 0.72 0.72

7

2 6.5

0 0 0 0 0 0 0 0 0

30 30

240 240 240 240 240 240

Catalyst No. 9

Catalyst No. 10

0.015

0.36

0.5

7 6 2 10 a 7

9

240 240

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240 240 240 240 240

30

0.72 0.65 0.65 0.50 0.50

6.0

0 0 0

> Catalyst No. 20 Catalyst No. 21

6.0 0.7 0.7

0.72

2

0.019 0.019 0.075

0.075

6.7

0.024

0.022

0.024

0.72

0.014 0.054

0.36 1.44

0.5

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Catalyst No. 12 Catalyst No. 13 Catalyst No. 14 Catalyst No. 15 Catalyst No. 16 Catalyst No. 17 Catalyst No. 18 Catalyst No. 19

Catalyst No. 11

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0.031

0.001	
2.99	
90'0	
1	
 70	
0	
30	
19.7	
34	

-																	
0.075	0.075	0.081	0.022	0:050	*Z/*N	0.000	0.013	0.013	0.099	0.046	0.000	0.007	0.007	0.099	0.026	0.013	*Z/*N
6.7	1.9	13.3	33.3	20.0	*Z	66.7	2.99	66.7	2.99	66.7	66.7	66.7	2.99	66.7	66.7	66.7	*Z
0.50	0.50	1.08	0.72	1.01	*z	00:0	0.88	0.88	6.61	3.08	0.00	0.44	0.44	19.9	1.76	0.88	*z
0.7	0.7	1.5	1	1.4	N- ppm	0	20	20	150	70	0	10	10	150	40	20	NO ppm
7	2	4	10	9	VCL ppm	70	20	20	20	20	20	20	20	20	20	20	VCL ppm
0	0	0	0	0	N- compounds	none	ON	NO2	NH3	C2H4(NH2)2	none	NO	NO2	NH3	C2H4(NH2)2	CH3CN	C2H6%
30	30	30	30	30	C2H4 (%)	30	30	30	30	30	30	30	30	30	30	30	C2H4 (%)
240	240	240	240	240	P (psia)	14.7	14.7	14.7	14.7	14.7	14.7	14.7	14.7	14.7	14.7	14.7	P (psia)
Catalyst No. 22	Catalyst No. 23	Catalyst No. 24	Catalyst No. 25	Catalyst No. 26	Example No. 15	Catalyst No. 31	Catalyst No. 32	Example No. 18									

-			 				
0.046	0.054	0.065	*Z/*N	0.018	0.018	0.018	0.018
23.3	20.0	16.7	*Z	33,3	33.3	33.3	33.3
1.08	1.08	1.08	* Z	0.59	0.59	0.59	0.59
1.5	1.5	1.5	mdd ON	10	10	10	10
7	9	5	VCL equivalent ppm	01	10	10	10
0	0	0	C2H6%	0	0	0	0
30	30	30	C2H4 (%)	30	30	30	30
240	240	240	P (psia)	19.7	19.7	19.7	19.7
Catalyst No. 35 (at 7 ppm vinyl chloride)	Catalyst No. 35 (at 6 ppm vinyl chloride)	Catalyst No. 35 (at 5 ppm vinyl chloride)	Example No. 22	Catalyst No. 38 (10 ppm vinyl chloride)	Catalyst No. 38 (10 ppm dichloroethane)	Catalyst No. 38 (50 ppm methyl chloride)	Catalyst No. 38 (4 ppm 2- chloropropane)